

Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L1	775901	array or arrays or microarray or microarrays	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/05/02 11:39
L2	29625	I1 and oligonucleotide\$	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/05/02 11:40
L3	27932	I2 and DNA	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/05/02 11:40
L4	211	I3 and 'different concentration'	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/05/02 11:40

FILE 'HOME' ENTERED AT 15:49:17 ON 02 MAY 2005

FILE 'CA' ENTERED AT 15:49:26 ON 02 MAY 2005
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=> s array? or microarray?
106403 ARRAY?
27634 MICROARRAY?
L1 127537 ARRAY? OR MICROARRAY?

=> s l1 and (dna or oligonucleotide?)
692986 DNA
72773 OLIGONUCLEOTIDE?
L2 28139 L1 AND (DNA OR OLIGONUCLEOTIDE?)

=> s l2 and (different concentration?)
1998743 DIFFERENT
203538 CONCENTRATION?
1195 DIFFERENT CONCENTRATION?
(DIFFERENT (W) CONCENTRATION?)
L3 0 L2 AND (DIFFERENT CONCENTRATION?)

=> s l2 and (different (8w) concentration?)
1998743 DIFFERENT
203538 CONCENTRATION?
2574 DIFFERENT (8W) CONCENTRATION?
L4 0 L2 AND (DIFFERENT (8W) CONCENTRATION?)

=> s 12 and concentration?
203538 CONCENTRATION?
L5 74 L2 AND CONCENTRATION?

```
=> s 15 and different  
          1998743 DIFFERENT  
L6          12 L5 AND DIFFERENT
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=> d ti ab 1-12

L6 ANSWER 1 OF 12 CA COPYRIGHT 2005 ACS on STN
TI Hybridization isotherms of **DNA microarrays** and the quantification of mutation studies
AB Diagnostic **DNA arrays** for detection of point mutations as markers for cancer usually function in the presence of a large excess of wild type **DNA**. This excess can give rise to false positives due to competitive hybridization of the wild type target at the mutation spot. The anal. of the **DNA array** data is typically qual. aiming to establish the presence or absence of a particular point mutation. Our theor. approach yields methods for quantifying the anal. so as to obtain the ratio of concns. of mutated and wild type **DNA**. The theory is formulated in terms of the hybridization isotherms relating the hybridization fraction at the spot to the composition of the sample solns at thermodn. equilibrium It focuses on samples containing an excess of single stranded **DNA** and on **DNA arrays** with low

surface d. of probes. The hybridization equilibrium consts. can be obtained by the nearest neighbor method. Two approaches allow us to obtain quant. results from the **DNA array** data. In one the signal of the mutation spot is compared with that of the wild type spot. The implementation requires knowledge of the saturation intensity of the two spots. The second approach requires comparison of the intensity of the mutation spot at two **different** temps. In this case knowledge of the saturation signal is not always necessary. **DNA arrays** can be used to obtain quant. results on the concentration ratio of mutated **DNA** to wild type **DNA** in studies of somatic point mutations.

- L6 ANSWER 2 OF 12 CA COPYRIGHT 2005 ACS on STN
TI Analytic chip for quantifying nucleic acid **concentration**, analytic device for quantifying nucleic acid **concentration** and analytic method for quantifying nucleic acid **concentration**
AB The invention relates to a device having a plural number of working electrodes, each carrying a single nucleic acid probe having a nucleic acid complementary with a target nucleic acid which is immobilized thereon and being **different** from each other in sensor area, and a normalization circuit for normalizing detection signals obtained in the working electrodes concerning resp. sensor areas.
- L6 ANSWER 3 OF 12 CA COPYRIGHT 2005 ACS on STN
TI Relationship between G+C content, ORF-length and mRNA **concentration** in *Saccharomyces cerevisiae*
AB RNA biogenesis is a tightly-regulated process. The levels and timing of expression of a gene depends on its particular function. However, gene expression levels may also depend on structural features. Here we describe the anal. of gene expression of 4977 ORFs using **DNA microarrays** covering the whole genome of three **different** *S. cerevisiae* strains, wild-type and tho2 and thp1 mutants with a general effect on mRNA biogenesis. We show that transcripts from G+C-rich ORFs accumulate at higher concns. than those from G+C-poor ones, in **different** ORF-length categories in all strains tested. In addition, we found a neg. correlation between ORF length and G+C content. Our results indicate that length and G+C content of a gene have a clear effect on its levels of expression. We discuss the biol. relevance of these results, as well as **different** ways that these structural features could modulate mRNA biogenesis.
- L6 ANSWER 4 OF 12 CA COPYRIGHT 2005 ACS on STN
TI Injector-concentrator electrodes for microchannel electrophoresis
AB An input port geometry, with injector-concentrator electrodes, for planar microchannel **array** for electrophoresis. This input port geometry enables efficient extraction and injection of the **DNA** sample from a single input port. The geometry, which utilizes injector-concentrator electrodes, allows simultaneous concentration, in **different** channels, of the sample into a longitudinally narrow strip just before releasing it for a run with enhanced injection spatial resolution, and time resolution. Optional multiple electrodes, at a **different** bias than the concentrator electrodes, may be used to discriminate against sample impurity ions. Electrode passivation can be utilized to prevent electrolysis. An addnl. electrode in or on the input hole can better define the initial loading. The injector-concentrator electrodes are positioned so that they cross the drift channel in a narrow strip at the bond plane between the top and bottom plates of the instrument and are located close to the inlet hole. The optional sample purification electrodes are located at a greater distance from the input hole than the injector-concentrate electrodes.
- L6 ANSWER 5 OF 12 CA COPYRIGHT 2005 ACS on STN
TI Transcriptional regulation and signature patterns revealed by **microarray** analyses of *Streptococcus pneumoniae* R6 challenged with

AB sublethal concentrations of translation inhibitors
The effects of sublethal concns. of four different classes of translation inhibitors (puromycin, tetracycline, chloramphenicol, and erythromycin) on global transcription patterns of *Streptococcus pneumoniae* R6 were determined by microarray analyses. Consistent with the general mode of action of these inhibitors, relative transcript levels of genes that encode ribosomal proteins and translation factors or that mediate tRNA charging and amino acid biosynthesis increased or decreased, resp. Transcription of the heat shock regulon was induced only by puromycin or streptomycin treatment, which lead to truncation or mistranslation, resp., but not by other antibiotics that block translation, transcription, or amino acid charging of tRNA. In contrast, relative transcript amts. of certain genes involved in transport, cellular processes, energy metabolism, and purine nucleotide (pur) biosynthesis were changed by different translation inhibitors. In particular, transcript amts. from a pur gene cluster and from purine uptake and salvage genes were significantly elevated by several translation inhibitors, but not by antibiotics that target other cellular processes. Northern blotting confirmed increased transcript amts. from part of the pur gene cluster in cells challenged by translation inhibitors and revealed the presence of a 10-kb transcript. Purine metabolism genes were neg. regulated by a homolog of the PurR regulatory protein, and full derepression in a ΔpurR mutant depended on optimal translation. Unexpectedly, hierarchical clustering of the microarray data distinguished among the global transcription patterns caused by antibiotics that inhibit different steps in the translation cycle. Together, these results show that there is extensive control of transcript amts. by translation in *S. pneumoniae*, especially for de novo purine nucleotide biosynthesis. In addition, these global transcription patterns form a signature that can be used to classify the mode of action and potential mechanism of new translation inhibitors.

L6 ANSWER 6 OF 12 CA COPYRIGHT 2005 ACS on STN
TI Microarrays and their manufacture by slicing bundled compound-containing fibers
AB Microarrays are prepared by using a sep. fiber for each compound being used in the microarray. The fibers are bundled and sectioned to form a thin microarray that may be glued to a backing. Antibodies to human serum albumin, transferrin, and haptoglobin were immobilized and crosslinked to Poros G particles. Each of the types of antibody-bearing particles plus particles free of antibodies was mixed with melted agarose and a different food coloring and sucked into a length of 1 mm diameter plastic tubing and gelled into rods. The rods were laid into an aluminum channel with more melted agarose to form an array of parallel rods embedded in a square cross-section bar of agarose. After the bar gelled, the gel was removed from the mold and transverse sections were prepared by slicing thin slices perpendicular to the axis of the bar and mounted on a glass slide.

L6 ANSWER 7 OF 12 CA COPYRIGHT 2005 ACS on STN
TI Extensions of counterion condensation theory. I. Alternative geometries and finite salt concentration
AB The counterion condensation theory originally proposed by Manning is extended to take account of both finite counterion concentration (mC) and the actual structure of the array of discrete charges. Counterion condensation is treated here as a binding isotherm problem, in which the unknown free volume is replaced by an unknown local binding constant β' , which is expected to vary with mC and polyion structure. The relation between the condensed fraction of counterion charge, r , β' and mC is obtained from the relevant grand partition function via the maximum term method. In the case of the single polyion in a large salt reservoir, the result is practically identical to Manning's equation. In order to determine the values of β' and r at arbitrary mC , a second relation between r , β' and mC is required. We propose an alternative auxiliary relation

that is equivalent to previous assumptions near $mC=0$, but which yields qual. correct and quant. useful results at finite mC . Simple expressions for r vs. mC and β' vs. mC are obtained by simultaneously solving the binding isotherm and auxiliary equations. Then r and β' are evaluated for five different linear arrays of infinite extent with different geometries: (1) a straight line of charges with uniform axial spacing; (2) two parallel lines of in-phase uniformly spaced charges; (3) a single-helix of discrete charges with uniform axial spacing; (4) a double-helix of discrete charges with uniform axial spacing of pairs of charges; (5) a cylindrical array of many parallel charged lines, chosen to simulate a uniformly charged cylinder. In all cases, the computed binding isotherms exhibit qual. correct behavior. As mC approaches zero, r approaches the Manning limit, $r=1-1/(LB/b)$ where b is the average axial spacing of electronic charges in the array and LB is the Bjerrum length. However, β' varies with polyion geometry, even in the zero salt limit, and matches the Manning value only in the case of a single straight charged line. With increasing mC , r declines significantly below its limiting value whenever $\lambda b > \sim 0.3$, where λ is the Debye screening parameter. In the case of cylindrical arrays containing either 2 or 100 parallel charged lines, r also decreases, whenever $\lambda d > \sim 2.0$, where d is the diameter of the array. In the case of two parallel charged lines, each with axial charge spacing $b=3.4$ Å, which are separated by $d=200$ Å, r exhibits a plateau value, 0.76, characteristic of the two combined lines, when $\lambda d < 2.0$, and declines with increasing mC to a shelf value, 0.52, characteristic of either single line, when $\lambda d > \sim 2.0$ and the lines become effectively screened from one another. β' Behaves in a roughly similar fashion. In the case of a cylindrical array of charged lines with the diameter and linear charge d of DNA, the r -values predicted by the present theory agree fairly well with those predicted by non-linear Poisson-Boltzmann theory up to 0.15 M uni-univalent salt.

L6 ANSWER 8 OF 12 CA COPYRIGHT 2005 ACS on STN
TI Biochip having probe molecules deposited in predetermined spatial concentration patterns
AB The present invention provides a method of and apparatus for manufacturing a biochip
in which different probe mols. are deposited on the substrate of the biochip with different concns. in accordance with a predetd. spatial concentration pattern. The present invention also provides a test apparatus
which uses the known concentration patterns to identify reactions of the different types of probe mols. with a test sample.

L6 ANSWER 9 OF 12 CA COPYRIGHT 2005 ACS on STN
TI Probing electrical properties of oriented DNA by conducting atomic force microscopy
AB Different methods have been applied for the stretching of DNA mols. on chemical functionalized surfaces by various modified reagents, i.e. 3-aminopropyltriethoxysilane or polylysine on mica and 2-mercaptopethylamine on Au(111)/mica by a moving interface technique, magnesium cation (Mg^{2+}) on mica by a spin-stretching method and DNA on an atomic-level flat mica by a free-flowing method. The long λ -DNA mol. is well elongated using the moving interface technique. The DNA mol. array d. can be controlled by the change of surface charge d. and the DNA concentration. On the other hand, the novel free-flowing method is very useful for the alignment of short polynucleotide mols. Shadow-mask evaporation has been used to fabricate a gold electrode contacted elec. to the oriented DNA mols. The intrinsic elec. properties of individual DNA mols. are directly measured using a conducting probe atomic force microscope equipped with a gold-coated conductive tip. The DNA mol. is considered as a promising mol. wire.

- L6 ANSWER 10 OF 12 CA COPYRIGHT 2005 ACS on STN
TI Apparatus and method to measure the activation state of signaling pathways in cells
AB The invention concerns the activity of multiple proteins in a single living cell, portion of a cell or in a group of cells simultaneously measured by introducing reporter mols. The reporter(s) is chemical modified by the enzyme of interest. In some cases the enzyme(s) is affected by the addition of a stimulus or a pharmaceutical compound to the cell. The reactions between the enzymes and the reporters are diminished or terminated, and the reporter and modified reporter are removed. The activity of the enzyme(s) is determined by measuring the amount of reporter remaining, the amount of altered reporter produced, or by comparing the amount of reporter to the amount of altered reporter. A database is compiled of the activities of the different proteins. By performing a series of expts. at different time points, conditions, and varieties of cell types, a database is developed for mol. cellular mechanisms in health and disease states. By exposing cells to variety of compds. data for drug development and screening is provided. Diagrams describing the apparatus assembly and operation are given.
- L6 ANSWER 11 OF 12 CA COPYRIGHT 2005 ACS on STN
TI Relationship of codon bias to mRNA concentration and protein length in *Saccharomyces cerevisiae*
AB In 1982, Ikemura reported a strikingly unequal usage of different synonymous codons, in five *Saccharomyces cerevisiae* nuclear genes having high protein levels. To study this trend in detail, we examined data from three independent studies that used oligonucleotide arrays or SAGE to estimate mRNA concns. for nearly all genes in the genome. Correlation coeffs. were calculated for the relationship of mRNA concentration to four commonly used measures of synonymous codon usage bias: the codon adaptation index (CAI), the codon bias index (CBI), the frequency of optimal codons (Fop), and the effective number of codons (Nc). mRNA concentration was best approximated as an exponential function of each of these four measures. Of the four, the CAI was the most strongly correlated with mRNA concentration ($rs=0.62\pm0.01$, $n=2525$, $p<10^{-17}$). When we controlled for CAI, mRNA concentration and protein length were neg. correlated (partial $rs=-0.23\pm0.01$, $n=4765$, $p<10^{-17}$). This may result from selection to reduce the size of abundant proteins to minimize transcriptional and translational costs. When we controlled for mRNA concentration, protein length and CAI were pos. correlated (partial $rs=0.16\pm0.01$, $n=4765$, $p<10^{-17}$). This may reflect more effective selection in longer genes against missense errors during translation. The correlation coeffs. between the mRNA levels of individual genes, as measured by different investigators and methods, were low, in the range $rs=0.39-0.68$.
- L6 ANSWER 12 OF 12 CA COPYRIGHT 2005 ACS on STN
TI Method and apparatus for detecting low concentrations of (bio)chemical components in a test medium using surface plasmon resonance
AB A method and an apparatus are described for detecting low concns. of ≥ 1 (bio)chemical component present in a test medium in a test cell, having a metal layer as sub wall with an external glass prism, using the surface plasmon resonance effect. A light ray is coupled in and, after attenuated total reflection, is coupled out and the intensity thereof is measured. The incidence angle position of the resonance curve is determined under the influence of the change, caused by the component, in the dielec. constant to the test medium near the metal layer. An adjustable selector is applied to the metal layer, in order to influence the incidence angle position of the resonance curve, through which the concns. or concentration changes of ≥ 1 components in the test medium can be simultaneously determined through ≥ 1 differential measurement. A preferential association and

therefore a higher concentration at the metal layer of 1 component above another

is brought about. The adjustable selector at the metal layer is formed by an **array** of (bio)chemical affinity ligands which are fixedly adsorbed to the metal layer. The **array** of (bio)chemical affinity ligands are a number of **different** antibodies, antigens, or **DNA**- or **RNA**-probes, such that in the test medium various antigens, antibodies, or homologous **DNA**, resp., can be determined

=>.d all 8

L6 ANSWER 8 OF 12 CA COPYRIGHT 2005 ACS on STN
AN 138:1933 CA
ED Entered STN: 26 Dec 2002
TI Biochip having probe molecules deposited in predetermined spatial **concentration** patterns
IN Jones, Aled Wynne; Beckett, Martin Gregory
PA Scientific Generics Limited, UK
SO PCT Int. Appl., 51 pp.
CODEN: PIXXD2
DT Patent
LA English
IC ICM B01J019-00
ICS B01L003-00
CC 9-1 (Biochemical Methods)

FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI WO 2002096552	A2	20021205	WO 2002-GB2567	20020605
WO 2002096552	A3	20030410		
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW			
RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			

PRAI GB 2001-13358 A 20010601

CLASS

PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
WO 2002096552	ICM	B01J019-00
	ICS	B01L003-00

AB The present invention provides a method of and apparatus for manufacturing a biochip

in which **different** probe mols. are deposited on the substrate of the biochip with **different** concns. in accordance with a predetd. spatial concentration pattern. The present invention also provides a test apparatus

which uses the known concentration patterns to identify reactions of the **different** types of probe mols. with a test sample.

ST biochip probe deposition concn pattern

IT CCD cameras

Computer program

Computers

Concentration (condition)

DNA microarray technology

Fluorometry

Immobilization, molecular or cellular

Microarray technology

Protein **microarray** technology
(biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT **DNA**
Proteins
RL: RCT (Reactant); RACT (Reactant or reagent)
(biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Gases
Gels
Liquids
(carrying probe onto substrate; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Information systems
(data; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Apparatus
(for biochip manufacturing and for testing biochips; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Electric field
(having predetd. spatial pattern; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT **DNA**
Proteins
RL: ARG (Analytical reagent use); DEV (Device component use); TEM (Technical or engineered material use); ANST (Analytical study); USES (Uses)
(immobilized, probe mols.; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Biosensors
(optical; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Chemicals
(probe mols.; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT **DNA** sequences
(probes for; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Cell
Eubacteria
Virus
(probes; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Information systems
(storage; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Bar code labels
(two-dimensional, on substrate surface; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Pressure
(waves having predetd. spatial pattern; biochip having probe mols. deposited in predetd. spatial concentration patterns)

=> e 'jones, aled wynne'/au
E1 1 JONES ZEBULON J R/AU
E2 2 JONES ZOE A/AU

=> s e169
L7 5 "JONES ALED WYNNE"/AU

=> d ti ab 1-5

- L7 ANSWER 1 OF 5 CA COPYRIGHT 2005 ACS on STN
TI Methods and apparatus for DNA sequencing
AB A carrier carries a computer program for base calling DNA bases from a dataset corresponding to observed traces obtained from electrophoresis. The program generates a database in the computer memory corresponding to a model of a DNA sequence and refines the model of the DNA sequence by the following operations making a change to the base sequence of the model, predicting the form of the traces from the modified model, comparing the predicted traces with the observed traces in dataset to generate a penalty function, determining whether or not to accept the modified model based on the value of the penalty function; and repeating operations until termination criteria have been achieved. The program has the advantage that it is less empirical and dependent on exptl. setup than existing programs such as Phred.
- L7 ANSWER 2 OF 5 CA COPYRIGHT 2005 ACS on STN
TI Biochip having probe molecules deposited in predetermined spatial concentration patterns
AB The present invention provides a method of and apparatus for manufacturing a biochip which in which different probe mols. are deposited on the substrate of the biochip with different concns. in accordance with a predetd. spatial concentration pattern. The present invention also provides a test apparatus which uses the known concentration patterns to identify reactions of the different types of probe mols. with a test sample.
- L7 ANSWER 3 OF 5 CA COPYRIGHT 2005 ACS on STN
TI Assay apparatus, assay method, and probe array for use in same
AB Assay apparatus is disclosed which comprises: probe means; a reaction volume for exposing said probe means to an analyte; and sensor means for detecting radiation emitted by said probe means in response to excitation; wherein said probe means comprises a plurality of probes arrayed in said reaction volume, and said sensor means and said reaction volume are coupled such that selective or simultaneous detection of radiation emitted from plural probes is permitted.
- L7 ANSWER 4 OF 5 CA COPYRIGHT 2005 ACS on STN
TI Method and apparatus for manufacture of biochip array
AB The invention concerns a method of and apparatus for manufacturing a biochip in which droplets containing at least one probe substance are deposited at random onto the biochip substrate. The probes are preferably sprayed using, for example, an aerosol nozzle or the like. In another embodiment, an electromagnetic, acoustic or optical deflector may be used to deflect the aerosol droplets in order to deposit the droplets onto the biochip in a pseudo-array. In a further embodiment, a test apparatus is provided in which a spatial intensity profile of a probe site is measured and used to reduce noise caused by, for example, scratches on the surface of the biochip. Diagrams describing the apparatus assembly and operation are given.
- L7 ANSWER 5 OF 5 CA COPYRIGHT 2005 ACS on STN
TI Sample processing apparatus
AB The present invention provides a system for processing biol. or chemical samples. The system includes a support for supporting the sample and a mount which is movable relative to the sample by a positioner. The positioner includes a number of flexure elements which are rigid along their axis of extent and flexible along the or each axis of extent of the other flexure elements. The flexure elements are rigidly secured at one of their ends to the mount and displaceably mounted at the other of their ends to an actuator which is operable to move the flexure element and hence move the mount.

=> d all 2

L7 ANSWER 2 OF 5 CA COPYRIGHT 2005 ACS on STN
AN 138:1933 CA
ED Entered STN: 26 Dec 2002
TI Biochip having probe molecules deposited in predetermined spatial concentration patterns
IN Jones, Aled Wynne; Beckett, Martin Gregory
PA Scientific Generics Limited, UK
SO PCT Int. Appl., 51 pp.
CODEN: PIXXD2
DT Patent
LA English
IC ICM B01J019-00
ICS B01L003-00
CC 9-1 (Biochemical Methods)
FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2002096552	A2	20021205	WO 2002-GB2567	20020605
	WO 2002096552	A3	20030410		
	W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW			
	RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			

PRAI GB 2001-13358 A 20010601

CLASS

PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
WO 2002096552	ICM	B01J019-00
	ICS	B01L003-00

AB The present invention provides a method of and apparatus for manufacturing a biochip

which in which different probe mols. are deposited on the substrate of the biochip with different concns. in accordance with a predetd. spatial concentration pattern. The present invention also provides a test apparatus which

uses the known concentration patterns to identify reactions of the different types of probe mols. with a test sample.

ST biochip probe deposition concn pattern

IT CCD cameras

Computer program

Computers

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Fluorometry

Immobilization, molecular or cellular

Microarray technology

Protein microarray technology

(biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT DNA

Proteins

RL: RCT (Reactant); RACT (Reactant or reagent)

(biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Gases

Gels
Liquids
 (carrying probe onto substrate; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Information systems
 (data; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Apparatus
 (for biochip manufacturing and for testing biochips; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Electric field
 (having predetd. spatial pattern; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT DNA
Proteins
RL: ARG (Analytical reagent use); DEV (Device component use); TEM (Technical or engineered material use); ANST (Analytical study); USES (Uses)
 (immobilized, probe mols.; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Biosensors
 (optical; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Chemicals
 (probe mols.; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT DNA sequences
 (probes for; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Cell
Eubacteria
Virus
 (probes; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Information systems
 (storage; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Bar code labels
 (two-dimensional, on substrate surface; biochip having probe mols. deposited in predetd. spatial concentration patterns)

IT Pressure
 (waves having predetd. spatial pattern; biochip having probe mols. deposited in predetd. spatial concentration patterns)

=> d all 3-5

L7 ANSWER 3 OF 5 CA COPYRIGHT 2005 ACS on STN
AN 137:165799 CA
ED Entered STN: 12 Sep 2002
TI Assay apparatus, assay method, and probe array for use in same
IN Laitenberger, Peter Georg; Disley, Darrin Matthew; Hember, Miles William
Noel; Jones, Aled Wynne
PA Scientific Generics Limited, UK
SO PCT Int. Appl., 70 pp.
CODEN: PIXXD2
DT Patent
LA English
IC ICM G01N021-64
CC 9-1 (Biochemical Methods)

FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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PI	WO 2002066965	A2	20020829	WO 2002-GB717	20020219
	WO 2002066965	A3	20030213		
	W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				

PRAI	GB 2001-4009	A	20010219
	GB 2001-4010	A	20010219

CLASS

PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
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WO 2002066965	ICM	G01N021-64

AB Assay apparatus is disclosed which comprises: probe means; a reaction volume for

exposing said probe means to an analyte; and sensor means for detecting radiation emitted by said probe means in response to excitation; wherein said probe means comprises a plurality of probes arrayed in said reaction volume, and said sensor means and said reaction volume are coupled such that selective or simultaneous detection of radiation emitted from plural probes is permitted.

ST array optical probe biomol detection

IT CCD cameras

Diffraction gratings

Electroluminescent devices

Fluorometry

High throughput screening

Interferometers

Lab-on-a-chip

Lasers

Lenses

Microarray technology

Micromachining

Optical fibers

Optical waveguides

Optics

Phosphorimetry

Sensors

Synchrotron radiation

(assay apparatus, assay method, and probe array for use in same)

IT DNA

RL: ANT (Analyte); ANST (Analytical study)

(assay apparatus, assay method, and probe array for use in same)

IT Borosilicates

RL: DEV (Device component use); USES (Uses)

(assay apparatus, assay method, and probe array for use in same)

L7 ANSWER 4 OF 5 CA COPYRIGHT 2005 ACS on STN

AN 137:151997 CA

ED Entered STN: 05 Sep 2002

TI Method and apparatus for manufacture of biochip array

IN Davies, Philip Andrew; Jones, Aled Wynne; Disley, Darrin Matthew; Hember, Miles William Noel; Miller, Nick; Hendry, Stuart Paul; Timson, Daniel Reginald Ewart

PA Scientific Generics Limited, UK

SO PCT Int. Appl., 61 pp.

CODEN: PIXXD2

DT Patent

LA English

IC ICM B01L003-02
ICS B01J019-00
CC 9-1 (Biochemical Methods)
Section cross-reference(s) : 1

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2002064255	A1	20020822	WO 2002-GB664	20020215
	W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM			
	RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
PRAI	GB 2001-3767	A	20010215		
	GB 2001-3768	A	20010215		

CLASS

PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
WO 2002064255	ICM	B01L003-02
	ICS	B01J019-00

AB The invention concerns a method of and apparatus for manufacturing a biochip in which

droplets containing at least one probe substance are deposited at random onto the biochip substrate. The probes are preferably sprayed using, for example, an aerosol nozzle or the like. In another embodiment, an electromagnetic, acoustic or optical deflector may be used to deflect the aerosol droplets in order to deposit the droplets onto the biochip in a pseudo-array. In a further embodiment, a test apparatus is provided in which a spatial intensity profile of a probe site is measured and used to reduce noise caused by, for example, scratches on the surface of the biochip. Diagrams describing the apparatus assembly and operation are given.

ST biochip array aerosol droplet protein DNA probe drug screening

IT Apparatus

(acoustic deflector; method and apparatus for manufacture of biochip array)

IT Information systems

(data, positional; method and apparatus for manufacture of biochip array)

IT Optical instruments

(deflectors; method and apparatus for manufacture of biochip array)

IT Apparatus

(electromagnetic deflector; method and apparatus for manufacture of biochip array)

IT Aerosols

Analytical apparatus

Animal cell

Biotechnology

Charge coupled devices

Chemicals

Computer program

Computers

Drug screening

Immobilization, molecular or cellular

Light

Liquids

Optical detectors

Storage

Virus

Water reservoirs

(method and apparatus for manufacture of biochip array)

IT DNA

Probes (nucleic acid)

Proteins

RL: ARG (Analytical reagent use); DEV (Device component use); ANST
(Analytical study); USES (Uses)

(method and apparatus for manufacture of biochip array)

IT Information, biological
(substrate serial number; method and apparatus for manufacture of biochip array)

RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

- (1) Commissariat Energie Atomique; FR 2791280 A 2000
- (2) Hahn Schickard Ges; DE 19913076 A 2000
- (3) Hahn Schickard Ges; DE 19947878 C 2001 CAPLUS
- (4) Tisone, T; US 6063339 A 2000 CA

L7 ANSWER 5 OF 5 CA COPYRIGHT 2005 ACS on STN

AN 137:87627 CA

ED Entered STN: 01 Aug 2002

TI Sample processing apparatus

IN Davies, Philip Andrew; Disley, Darrin Matthew; Jones, Aled Wynne
; Purvis, Duncan Ross; Beckett, Martin Gregory; Miller, Nicholas;
Miller-Jones, David Nicholas

PA Scientific Generics Limited, UK

SO PCT Int. Appl., 50 pp.

CODEN: PIXXD2

DT Patent

LA English

IC ICM B01L009-06

ICS G01N001-31; B01J019-00; G01N035-10; G01N035-04

CC 80-2 (Organic Analytical Chemistry)
Section cross-reference(s): 9

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2002055201	A2	20020718	WO 2002-GB122	20020114
	WO 2002055201	A3	20021114		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM					
	RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
PRAI	GB 2001-901	A	20010112		
	GB 2001-1896	A	20010124		
	GB 2001-1898	A	20010124		
	GB 2001-2344	A	20010130		

CLASS

PATENT NO. CLASS PATENT FAMILY CLASSIFICATION CODES

	WO 2002055201	ICM	B01L009-06	ICS G01N001-31; B01J019-00; G01N035-10; G01N035-04
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AB The present invention provides a system for processing biol. or chemical samples. The system includes a support for supporting the sample and a mount which is movable relative to the sample by a positioner. The positioner includes a number of flexure elements which are rigid along their axis of extent and flexible along the or each axis of extent of the other flexure elements. The flexure elements are rigidly secured at one of their ends to the mount and displaceably mounted at the other of their ends to an actuator which is operable to move the flexure element and hence move the mount.

ST processing app materials handling
IT Materials handling
 (apparatus; sample processing apparatus for biol. or chemical samples)
IT Actuators
 Bending
 Biological materials
 Microarray technology
 Optical fibers
 (sample processing apparatus for biol. or chemical samples)

=> s e143-e144

 1 "BECKETT MARTIN G"/AU
 2 "BECKETT MARTIN GREGORY"/AU
L8 3 ("BECKETT MARTIN G"/AU OR "BECKETT MARTIN GREGORY"/AU)

=> d ti ab 1-3

L8 ANSWER 1 OF 3 CA COPYRIGHT 2005 ACS on STN

TI Biochip having probe molecules deposited in predetermined spatial concentration patterns

AB The present invention provides a method of and apparatus for manufacturing a biochip
 in which different probe mols. are deposited on the substrate of the biochip with different concns. in accordance with a predetd. spatial concentration pattern. The present invention also provides a test apparatus which
 uses the known concentration patterns to identify reactions of the different types of probe mols. with a test sample.

L8 ANSWER 2 OF 3 CA COPYRIGHT 2005 ACS on STN

TI Sample processing apparatus

AB The present invention provides a system for processing biol. or chemical samples. The system includes a support for supporting the sample and a mount which is movable relative to the sample by a positioner. The positioner includes a number of flexure elements which are rigid along their axis of extent and flexible along the or each axis of extent of the other flexure elements. The flexure elements are rigidly secured at one of their ends to the mount and displaceably mounted at the other of their ends to an actuator which is operable to move the flexure element and hence move the mount.

L8 ANSWER 3 OF 3 CA COPYRIGHT 2005 ACS on STN

TI Infrared observations of gravitational lensing in Abell 2219 with CIRSI

AB We present the 1st detection of a gravitational depletion signal at near-IR wavelengths, based on deep panoramic images of the cluster Abell 2219 ($z = 0.22$) taken with the Cambridge IR survey instrument (CIRSI) at the prime focus of the 4.2-m William Herschel telescope. IR studies of gravitational depletion offer a number of advantages over similar techniques applied at optical wavelengths, and can provide reliable total masses for intermediate-red shift clusters. Using the maximum-likelihood technique developed by Schneider, King & Erben, we detect the gravitational depletion at the 3σ confidence level. By modeling the mass distribution as a singular isothermal sphere and ignoring the uncertainty in the unlensed number counts, we find an Einstein radius of $\theta_E \approx 13.7 - 4.2 + 3.9$ arcsec (66% confidence limit). This corresponds to a projected velocity dispersion of $\sigma_v \approx 800$ km s $^{-1}$, in agreement with constraints from strongly lensed features. For a Navarro, Frenk, & White mass model, the radial dependence observed indicates a best-fitting halo scale-length of 125 h $^{-1}$ kpc. We investigate the uncertainties arising from the observed fluctuations in the unlensed number counts, and show that clustering is the dominant source of error. We extend the maximum-likelihood method to include the effect of incompleteness, and discuss the prospects of further systematic studies of lensing in the

near-IR band.

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=> e 'shalon'/au
E1      2      SHALOMOVA T P/AU
E2      1      SHALOMYANSKY A M/AU
E3      0 ---> SHALON/AU
E4      1      SHALON D/AU
E5      7      SHALON DARI/AU
E6      6      SHALON R/AU
E7      1      SHALON RAHEL/AU
E8      1      SHALON S/AU
E9      9      SHALON TADMOR/AU
E10     1      SHALON TIDHAR D/AU
E11     3      SHALON TIDHAR DARI/AU
E12     2      SHALON Y/AU

=> s e10-e11
      1 "SHALON TIDHAR D"/AU
      3 "SHALON TIDHAR DARI"/AU
L9      4 ("SHALON TIDHAR D"/AU OR "SHALON TIDHAR DARI"/AU)

=> d ti ab 1-4
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L9 ANSWER 1 OF 4 CA COPYRIGHT 2005 ACS on STN
TI Capillary printing systems
AB The invention provides printing systems and methods for depositing small vols. of liquid on solid substrates and is particularly suited for printing high-d. anal. arrays. These systems and methods are useful with a wide variety of liqs. and substrates and offer a wide variety of applications, including the deposition of arrays of analytes. In particular embodiments, the systems comprise a preservation device, a detachable ganged plurality of printing devices, and/or a wire bonding capillary.

L9 ANSWER 2 OF 4 CA COPYRIGHT 2005 ACS on STN
TI Methods for fabricating microarrays of biological samples
AB A method and apparatus for forming microarrays of biol. samples on a support are disclosed. The method involves dispensing a known volume of a reagent at each selected array position, by tapping a capillary dispenser on the support under conditions effective to draw a defined volume of liquid onto the support. The apparatus is designed to produce a microarray of such regions in an automated fashion. The apparatus is used for genetic methods, e.g. microarray hybridization for gene expression with high partial concentration of each cDNA species; multiplex colorimetric hybridization on a gridded support; genomic complexity hybridization to DNA where microarrays represent the *Saccharomyces cerevisiae* genome etc.

L9 ANSWER 3 OF 4 CA COPYRIGHT 2005 ACS on STN
TI Dna micro arrays: a new tool for genetic analysis
AB Unavailable

L9 ANSWER 4 OF 4 CA COPYRIGHT 2005 ACS on STN
TI Method and apparatus for fabricating microarrays of biological samples
AB A method and apparatus for forming microarrays of biol. samples on a support are disclosed for, e.g., large-scale screening assays, such as arrays of DNA samples to be used in DNA hybridization assays for genetic research and diagnostic applications. The method involves dispensing a known volume of a reagent at each of a selected array position, by tapping a capillary dispenser on the support under conditions effective to draw a defined volume of liquid onto the support. The apparatus is designed to produce a microarray of such regions in an automated fashion.

=> d all 1-4

L9 ANSWER 1 OF 4 CA COPYRIGHT 2005 ACS on STN
AN 132:201091 CA
ED Entered STN: 31 Mar 2000
TI Capillary printing systems
IN Shalon, Tidhar D.; Maurino, Joseph R.; Titsworth, Loren D.;
Bevirt, Joeben
PA Incyte Pharmaceuticals, Inc., USA
SO PCT Int. Appl., 30 pp.
CODEN: PIXXD2
DT Patent
LA English
IC ICM B01L003-02
CC 74-6 (Radiation Chemistry, Photochemistry, and Photographic and Other
Reprographic Processes)
Section cross-reference(s): 3, 79, 80

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2000013796	A1	20000316	WO 1999-US20692	19990909
	W: AU, CA, JP				
	RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
	US 6309891	B1	20011030	US 1998-150502	19980909
	CA 2342785	AA	20000316	CA 1999-2342785	19990909
	AU 9959153	A1	20000327	AU 1999-59153	19990909
	AU 748153	B2	20020530		
	EP 1109624	A1	20010627	EP 1999-946833	19990909
	EP 1109624	B1	20040218		
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI				
	JP 2003519360	T2	20030617	JP 2000-568592	19990909
	EP 1374998	A1	20040102	EP 2003-20699	19990909
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI, CY				
	AT 259678	E	20040315	AT 1999-946833	19990909
	US 2001013298	A1	20010816	US 2001-819166	20010327
	US 2002064887	A1	20020530	US 2001-819172	20010327
	US 2001044157	A1	20011122	US 2001-884506	20010614
	JP 2004155201	A2	20040603	JP 2003-387270	20031117
PRAI	US 1998-150502	A	19980909		
	EP 1999-946833	A3	19990909		
	JP 2000-568592	A3	19990909		
	WO 1999-US20692	W	19990909		

CLASS

PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
WO 2000013796	ICM	B01L003-02
WO 2000013796	ECLA	B01J019/00C; B01L003/00C2D
US 6309891	NCL	436/180.000; 073/864.010; 101/494.000; 141/031.000; 422/100.000; 436/049.000; 436/054.000; 436/176.000
	ECLA	B01J019/00C; B01L003/00C2D; B01L003/02D
EP 1374998	ECLA	B01L003/02D
US 2001013298	NCL	101/494.000
	ECLA	B01J019/00C; B01L003/00C2D; B01L003/02D
US 2002064887	NCL	436/180.000; 422/100.000; 347/040.000
	ECLA	B01J019/00C; B01L003/00C2D; B01L003/02D
US 2001044157	NCL	436/180.000; 422/100.000; 422/131.000
	ECLA	B01J019/00C; B01L003/00C2D; B01L003/02D
JP 2004155201	FTERM	2C064/CC07; 2C064/CC08; 2C064/CC13; 4F041/AA02; 4F041/AB01; 4F041/BA02; 4F041/BA10; 4F041/BA12; 4F041/BA13; 4F041/BA36

AB The invention provides printing systems and methods for depositing small

vols. of liquid on solid substrates and is particularly suited for printing high-d. anal. arrays. These systems and methods are useful with a wide variety of liqs. and substrates and offer a wide variety of applications, including the deposition of arrays of analytes. In particular embodiments, the systems comprise a preservation device, a detachable ganged plurality of printing devices, and/or a wire bonding capillary.

ST Capillary printing system analysis array

IT Capillary vessel
Printing (nonimpact)
(capillary printing systems for depositing small vols. of liquid on solid substrates for printing high-d. anal. arrays in relation to)

IT Polynucleotides
RL: ARU (Analytical role, unclassified); ANST (Analytical study)
(capillary printing systems for depositing small vols. of liquid on solid substrates for printing high-d. anal. arrays in relation to)

IT Materials handling
(delivery apparatus; capillary printing systems for depositing small vols. of liquid on solid substrates for printing high-d. anal. arrays in relation to)

IT Samples
(liquid; capillary printing systems for depositing small vols. of liquid on solid substrates for printing high-d. anal. arrays in relation to)

RE.CNT 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

(1) Akihiro, K; US 5607861 A 1997 CA
(2) Hrubesh, C; US 4441532 A 1984
(3) Konrad, B; US 5763278 A 1998
(4) Roach, D; US 5770151 A 1998 CA
(5) Smith, M; US 4142656 A 1979
(6) Sohrab, D; WO 8910192 A 1989
(7) Univ Leland Stanford Junior; WO 9535505 A 1995 CA

L9 ANSWER 2 OF 4 CA COPYRIGHT 2005 ACS on STN
AN 129:226623 CA
ED Entered STN: 24 Oct 1998
TI Methods for fabricating microarrays of biological samples
IN Brown, Patrick O.; Shalon, Tidhar Dari
PA The Board of Trustees of the Leland Stanford Junior University, USA
SO U.S., 18 pp., Cont.-in-part of U.S. Ser. No. 261,388, abandoned.
CODEN: USXXAM
DT Patent
LA English
IC ICM C12M001-34
ICS C12M001-40
INCL 422050000
CC 3-1 (Biochemical Genetics)
FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 5807522 CA 2192095 CA 2192095 WO 9535505	A AA C A1	19980915 19951228 19990831 19951228	US 1995-477809 CA 1995-2192095 WO 1995-US7659	19950607 19950616 19950616
	W: AU, CA, JP RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE AU 9528629 AU 709276 EP 804731 EP 804731				
		A1	19960115	AU 1995-28629	19950616
		B2	19990826		
		A1	19971105	EP 1995-923921	19950616
		B1	19990526		
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE JP 10503841 EP 913485 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE AT 180570				
		T2	19980407	JP 1995-502498	19950616
		A1	19990506	EP 1998-123001	19950616
		E	19990615	AT 1995-923921	19950616

ES	2134481	T3	19991001	ES	1995-923921	19950616
JP	3272365	B2	20020408	JP	1996-502498	19950616
JP	2002243736	A2	20020828	JP	2001-132630	19950616
US	6110426	A	20000829	US	1997-1027	19971230
US	2003012695	A1	20030116	US	1998-356322	19981124
US	2001051344	A1	20011213	US	2001-908304	20010717
PRAI	US 1994-261388	B2	19940617			
	US 1995-477809	A	19950607			
	EP 1995-923921	A3	19950616			
	JP 1996-502498	A3	19950616			
	WO 1995-US7659	W	19950616			
	US 1995-514875	A2	19950814			
	US 1996-688488	A1	19960730			
	US 1998-188931	A1	19981110			

CLASS

PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
US 5807522	ICM	C12M001-34
	ICS	C12M001-40
	INCL	422050000
US 5807522	NCL	422/050.000; 422/052.000; 422/055.000; 422/056.000; 422/057.000; 422/068.100; 422/069.000; 422/082.050; 422/082.060; 422/082.070; 422/082.080; 435/006.000; 435/007.100; 436/501.000; 530/300.000; 530/333.000; 530/334.000; 530/350.000; 536/025.300
	ECLA	B01J019/00C; B01L003/02D; C12Q001/68A6; C12Q001/68B10A; G01N033/543K
WO 9535505	ECLA	B01J019/00C; B01L003/02D; C12Q001/68A6; C12Q001/68B10A; G01N033/543K
EP 913485	ECLA	B01J004/02; B01L003/02D; B01J019/00C
US 6110426	NCL	422/068.100; 422/050.000; 435/006.000; 435/283.100; 436/051.000; 536/025.300
	ECLA	B01J019/00C
US 2003012695	NCL	422/068.100; 435/006.000; 536/023.100
	ECLA	B01J004/02; B01J019/00C; B01L003/02D; C12Q001/68A6; C12Q001/68B10A; G01N033/543K
US 2001051344	NCL	435/006.000; 435/069.100; 422/068.100
	ECLA	B01L003/02D; C12Q001/68A6+565/501; C12Q001/68B10A+565/102

AB A method and apparatus for forming microarrays of biol. samples on a support are disclosed. The method involves dispensing a known volume of a reagent at each selected array position, by tapping a capillary dispenser on the support under conditions effective to draw a defined volume of liquid onto the support. The apparatus is designed to produce a microarray of such regions in an automated fashion. The apparatus is used for genetic methods, e.g. microarray hybridization for gene expression with high partial concentration of each cDNA species; multiplex colorimetric hybridization on a gridded support; genomic complexity hybridization to DNA where microarrays represent the *Saccharomyces cerevisiae* genome etc.

ST dispensing app capillary microarray hybridization DNA

IT Dispensing apparatus
(dosing; methods for fabricating microarrays of biol. samples)

IT Biological materials

Colorimetry

Genetic methods

Genome

Micromachining

Nucleic acid hybridization

Process automation

Saccharomyces cerevisiae

(methods for fabricating microarrays of biol. samples)

IT Reagents

RL: ARG (Analytical reagent use); ARU (Analytical role, unclassified);
ANST (Analytical study); USES (Uses)

(methods for fabricating microarrays of biol. samples)

IT DNA
CDNA

RL: BSU (Biological study, unclassified); BIOL (Biological study)
(methods for fabricating microarrays of biol. samples)

RE.CNT 70 THERE ARE 70 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

(1) Anon; WO 9003382 1990 CA
(2) Anon; WO 9210588 1992 CA
(3) Anon; WO 9322680 1993 CA
(4) Anon; WO 9500530 1995 CA
(5) Anon; WO 9515970 1995 CA
(6) Anon; WO 9521944 1995 CA
(7) Anon; WO 9525116 1995 CA
(8) Anon; EP 721016 A2 1996 CA
(9) Anon; WO 9617958 1996 CA
(10) Augenlicht; US 4981783 1991 CA
(11) Barrett; US 5252743 1993 CA
(12) Billings; FASEB 1991, V5, P28 CA
(13) Brennan; US 5474796 1995 CA
(14) Chang; US 4591570 1986 CA
(15) Chang; US 5100777 1992 CA
(16) Chateau; US 4071315 1978 CA
(17) Chee; Science 1996, V274, P610 CA
(18) Chiswell; US 4716106 1987 CA
(19) Cozzette; US 5200051 1993 CA
(20) Dattagupta; US 4670380 1987 CA
(21) Dattagupta; US 5348855 1994 CA
(22) Deeg; US 5338688 1994
(23) Douglas; US 5556748 1996 CA
(24) Drmanac; US 5202231 1993 CA
(25) Drmanac; DNA and Cell Biology 1990, V9, P527 CA
(26) Drmanac; Electrophoresis 1992, V13, P566 CA
(27) Drmanac; Science 1993, V260, P1649 CA
(28) Ekins; J Clinical Immunoassay 1990, V13(4), P169
(29) Fodor; US 5445934 1995 CA
(30) Fodor; US 5510270 1996 CA
(31) Gebeyehu; US 4921805 1990 CA
(32) Gilham; US 3730844 1973 CA
(33) Gong; US 5512430 1996 CA
(34) Grossman; US 5514543 1996 CA
(35) Heller; US 5605662 1997 CA
(36) Holmes; US 5242974 1993 CA
(37) Hozier; US 5563060 1996
(38) Ishii; US 5474895 1995 CA
(39) Kurn; US 4868104 1989 CA
(40) Lockhart; US 5556752 1996 CA
(41) Mathies; US 5091652 1992 CA
(42) Matsumoto; US 5204268 1993 CA
(43) McGall; US 5412087 1995 CA
(44) Mills; US 5064754 1991 CA
(45) Mullis; US 4683195 1987 CA
(46) Mullis; US 4683202 1987 CA
(47) Okano; US 5434049 1995 CA
(48) Oprandy; US 5200312 1993 CA
(49) Paau; US 4556643 1985 CA
(50) Palva; US 4731325 1988 CA
(51) Peters; US 5013669 1991
(52) Pirrung; US 5143854 1992 CA
(53) Rabbani; US 4755458 1988 CA
(54) Ranki; US 4486539 1984 CA
(55) Ranki; US 4563419 1986 CA
(56) Rava; US 5545531 1996 CA
(57) Sninsky; US 5389512 1995 CA

- (58) Soini; US 5028545 1991 CA
- (59) Soini; US 5518883 1996
- (60) Stapleton; US 5188963 1993 CA
- (61) Stokke; US 5472842 1995 CA
- (62) Trulson; US 5578832 1996 CA
- (63) Ullman; US 5185243 1993 CA
- (64) Ullman; US 5516641 1996 CA
- (65) Urdea; US 4868105 1989 CA
- (66) van Ness; US 5514785 1996 CA
- (67) Wallace; US 4767700 1988 CA
- (68) Ward; US 5328824 1994 CA
- (69) White; US 4677054 1987 CA
- (70) Zuckerman; US 5252296 1993

L9 ANSWER 3 OF 4 CA COPYRIGHT 2005 ACS on STN
 AN 124:308810 CA
 ED Entered STN: 29 May 1996
 TI Dna micro arrays: a new tool for genetic analysis
 AU **Shalon, Tidhar Dari**
 CS Stanford Univ., Stanford, CA, USA
 SO (1996) 108 pp. Avail.: Univ. Microfilms Int., Order No. DA9612036
 From: Diss. Abstr. Int., B 1996, 56(12), 6534
 DT Dissertation
 LA English
 CC 3-1 (Biochemical Genetics)
 Section cross-reference(s): 10, 11
 AB Unavailable
 ST DNA micro array genetics Arabidopsis Saccharomyces
 IT Arabidopsis thaliana
 Genetic mapping
 Genetics
 Saccharomyces cerevisiae
 (DNA micro arrays: a new tool for genetic anal. of gene expression and phys. mapping in organisms such as Arabidopsis and Saccharomyces)
 IT Gene
 RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)
 (expression, DNA micro arrays: a new tool for genetic anal. of gene expression and phys. mapping in organisms such as Arabidopsis and Saccharomyces)

L9 ANSWER 4 OF 4 CA COPYRIGHT 2005 ACS on STN
 AN 124:137802 CA
 ED Entered STN: 06 Mar 1996
 TI Method and apparatus for fabricating microarrays of biological samples
 IN **Shalon, Tidhar Dari**; Brown, Patrick O.
 PA Board of Trustees of the Leland Stanford Junior University, USA
 SO PCT Int. Appl., 52 pp.
 CODEN: PIXXD2

DT Patent
 LA English
 IC ICM G01N033-543
 ICS G01N033-68
 CC 3-1 (Biochemical Genetics)
 Section cross-reference(s): 9

FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 9535505	A1	19951228	WO 1995-US7659	19950616
	W: AU, CA, JP				
	RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
	US 5807522	A	19980915	US 1995-477809	19950607
	AU 9528629	A1	19960115	AU 1995-28629	19950616
	AU 709276	B2	19990826		

EP 804731	A1	19971105	EP 1995-923921	19950616
EP 804731	B1	19990526		
	R:	AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE		
JP 10503841	T2	19980407	JP 1995-502498	19950616
PRAI US 1994-261388	A	19940617		
US 1995-477809	A	19950607		
WO 1995-US7659	W	19950616		

CLASS

PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
-----	-----	-----
WO 9535505	ICM	G01N033-543
	ICS	G01N033-68
WO 9535505	ECLA	B01J019/00C; B01L003/02D; C12Q001/68A6; C12Q001/68B10A; G01N033/543K
US 5807522	NCL	422/050.000; 422/052.000; 422/055.000; 422/056.000; 422/057.000; 422/068.100; 422/069.000; 422/082.050; 422/082.060; 422/082.070; 422/082.080; 435/006.000; 435/007.100; 436/501.000; 530/300.000; 530/333.000; 530/334.000; 530/350.000; 536/025.300
	ECLA	B01J019/00C; B01L003/02D; C12Q001/68A6; C12Q001/68B10A; G01N033/543K

AB A method and apparatus for forming microarrays of biol. samples on a support are disclosed for, e.g., large-scale screening assays, such as arrays of DNA samples to be used in DNA hybridization assays for genetic research and diagnostic applications. The method involves dispensing a known volume of a reagent at each of a selected array position, by tapping a capillary dispenser on the support under conditions effective to draw a defined volume of liquid onto the support. The apparatus is designed to produce a microarray of such regions in an automated fashion.

ST biol sample automated reagent dispensing app; DNA hybridization assay
microarray prepn app; gene expression hybridization assay app

IT Dispensing apparatus
(automatic; method and apparatus for fabricating microarrays of biol. samples)

IT Samples
(biol.; method and apparatus for fabricating microarrays of biol. samples)

IT Gene
RL: ANT (Analyte); ANST (Analytical study)
(expression; method and apparatus for fabricating microarrays of biol. samples)

IT Analysis
Arabidopsis
Genome
Holders
Immobilization, biochemical
Nucleic acid hybridization
Plant analysis
Polymer-supported reagents
Polymerase chain reaction
Saccharomyces cerevisiae
(method and apparatus for fabricating microarrays of biol. samples)

IT Biopolymers
Deoxyribonucleic acids
Nucleic acids
Peptides, analysis
RL: ANT (Analyte); ANST (Analytical study)
(method and apparatus for fabricating microarrays of biol. samples)

IT Named reagents and solutions
RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)
(method and apparatus for fabricating microarrays of biol. samples)

IT Glass, oxide
RL: ARU (Analytical role, unclassified); DEV (Device component use); ANST (Analytical study); USES (Uses)

(slides; method and apparatus for fabricating microarrays of biol. samples)
IT Deoxyribonucleic acids
RL: ANT (Analyte); ANST (Analytical study)
(complementary, method and apparatus for fabricating microarrays of biol.
samples)
IT Nucleotides, analysis
RL: ANT (Analyte); ANST (Analytical study)
(poly-, method and apparatus for fabricating microarrays of biol. samples)
IT Chemicals
(reagents, method and apparatus for fabricating microarrays of biol.
samples)
IT 2321-07-5D, Fluorescein, oligonucleotides labeled with
RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)
(method and apparatus for fabricating microarrays of biol. samples)
IT 9004-70-0, Nitrocellulose 25104-18-1, Polylysine 38000-06-5,
Polylysine
RL: ARU (Analytical role, unclassified); DEV (Device component use); ANST
(Analytical study); USES (Uses)
(method and apparatus for fabricating microarrays of biol. samples)

=> d his

(FILE 'HOME' ENTERED AT 15:49:17 ON 02 MAY 2005)

FILE 'CA' ENTERED AT 15:49:26 ON 02 MAY 2005

L1 127537 S ARRAY? OR MICROARRAY?
L2 28139 S L1 AND (DNA OR OLIGONUCLEOTIDE?)
L3 0 S L2 AND (DIFFERENT CONCENTRATION?)
L4 0 S L2 AND (DIFFERENT(8W)CONCENTRATION?)
L5 74 S L2 AND CONCENTRATION?
L6 12 S L5 AND DIFFERENT
E 'JONES, ALED WYNNE'/AU
E 'JONES, ALED'/AU
E 'JONES, A?'/AU
E 'JONES'/AU
L7 5 S E169
E 'BECKETT, MARTIN'/AU
E 'BECKETT'/AU
L8 3 S E143-E144
E 'SHALON'/AU
L9 4 S E10-E11

=> s 12 and (concentration(10w)spot?)

140649 CONCENTRATION
96294 SPOT?
57 CONCENTRATION(10W)SPOT?
L10 0 L2 AND (CONCENTRATION(10W)SPOT?)

=> s 15 not 16

L11 62 L5 NOT L6

=> d his

(FILE 'HOME' ENTERED AT 15:49:17 ON 02 MAY 2005)

FILE 'CA' ENTERED AT 15:49:26 ON 02 MAY 2005
L1 127537 S ARRAY? OR MICROARRAY?
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	E 'JONES, ALED' /AU
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	E 'JONES' /AU
L7	5 S E169
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	E 'BECKETT' /AU
L8	3 S E143-E144
	E 'SHALON' /AU
L9	4 S E10-E11
L10	0 S L2 AND (CONCENTRATION(10W) SPOT?)
L11	62 S L5 NOT L6

=> b biosis

COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	231.75	231.96
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL SESSION
CA SUBSCRIBER PRICE	-63.92	-63.92

FILE 'BIOSIS' ENTERED AT 16:22:27 ON 02 MAY 2005
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FILE COVERS 1969 TO DATE.

CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNS) PRESENT
 FROM JANUARY 1969 TO DATE.

RECORDS LAST ADDED: 27 April 2005 (20050427/ED)

FILE RELOADED: 19 October 2003.

=> s 12

42543 ARRAY?
15873 MICROARRAY?
1068260 DNA
48348 OLIGONUCLEOTIDE?
L12 16406 L1 AND (DNA OR OLIGONUCLEOTIDE?)

=> s 112 not 12

42543 ARRAY?
15873 MICROARRAY?
1068260 DNA
48348 OLIGONUCLEOTIDE?
L13 0 L12 NOT L2

=> b medline

COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	0.85	232.81
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL SESSION
CA SUBSCRIBER PRICE	0.00	-63.92

FILE 'MEDLINE' ENTERED AT 16:22:54 ON 02 MAY 2005

FILE LAST UPDATED: 30 APR 2005 (20050430/UP). FILE COVERS 1950 TO DATE.

On December 19, 2004, the 2005 MeSH terms were loaded.

The MEDLINE reload for 2005 is now available. For details enter HELP RLOAD at an arrow prompt (=>). See also:

<http://www.nlm.nih.gov/mesh/>
http://www.nlm.nih.gov/pubs/techbull/nd04/nd04_mesh.html

OLDMEDLINE now back to 1950.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2005 vocabulary.

This file contains CAS Registry Numbers for easy and accurate substance identification.

```
=> s 12
    48736 ARRAY?
    12430 MICROARRAY?
    797644 DNA
    67604 OLIGONUCLEOTIDE?
L14      20445 L1 AND (DNA OR OLIGONUCLEOTIDE?)

=> s l14 not 12
    48736 ARRAY?
    12430 MICROARRAY?
    797644 DNA
    67604 OLIGONUCLEOTIDE?
L15      0 L14 NOT L2

=> s l14 and different (w) concentration?
    1068128 DIFFERENT
    1037046 CONCENTRATION?
    9945 DIFFERENT (W) CONCENTRATION?
L16      17 L14 AND DIFFERENT (W) CONCENTRATION?

=> d ti ab 1-17

=> d all 10

L16  ANSWER 10 OF 17      MEDLINE on STN
AN  2003502771      MEDLINE
DN  PubMed ID: 14579528
TI  Validation of cDNA microarray technology.
AU  Luo Yao; Xu Hong; Li Yao; Han Zhi-Yong; Qiu Min-Yan; Chen Qin; Liu
    San-Zhen; Ni Sheng; Xie Yi; Mao Yu-Min
CS  State Key Laboratory of Genetic Engineering, Institute of Genetics, School
    of Life Science, Fudan University, Shanghai 200433, China..
    yao_luo@hotmail.com
SO  Yi chuan xue bao = Acta genetica Sinica, (2003 Jul) 30 (7) 611-8.
    Journal code: 7900784. ISSN: 0379-4172.
CY  China
DT  Journal; Article; (JOURNAL ARTICLE)
    (VALIDATION STUDIES)
LA  Chinese
FS  Priority Journals
EM  200312
ED  Entered STN: 20031029
    Last Updated on STN: 20031219
    Entered Medline: 20031211
AB  cDNA microarray is a technological approach that has the
    potential to globally measure changes in mRNA expression levels.
    Self-comparison experiments with the same kind of tissue and differential
    expression experiments with the different kinds of tissue have been done
    to verify the reproducibility and the accuracy of this technique. The
    parameter of the reliability and the reproducibility of the
    microarray data were analyzed by correlation coefficient (R),
    coefficient of variation (CV) and false positive rate (FPR) etc.
    Meanwhile, the error resource also has been inspected. These results
```

showed that generally the correlation coefficient of data from this cDNA **microarray** system was more than 0.9, the coefficient of variation was about 15%, and the false positive rate was below 3%. The result proves the accuracy of the cDNA **microarray** data. Consistence rate (CR) was advanced here as a new parameter to evaluate the reproducibility of two replicate experiments. It has some advantages over correlation coefficient and coefficient of variation. The influence of some important factors in the experiments, such as **different concentration** of spotted **DNA**, mRNA and total RNA, different batches of slides and different processes of labeling, have been investigated by comparing the results. It was shown that most of the false position produced by the experiment system could be reduced by replicate experiments.

CT English Abstract
Humans
*Oligonucleotide Array Sequence Analysis: MT, methods
Reproducibility of Results

=> d his

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E 'JONES'/AU
L7 5 S E169
E 'BECKETT, MARTIN'/AU
E 'BECKETT'/AU
L8 3 S E143-E144
E 'SHALON'/AU
L9 4 S E10-E11
L10 0 S L2 AND (CONCENTRATION(10W) SPOT?)
L11 62 S L5 NOT L6

FILE 'BIOSIS' ENTERED AT 16:22:27 ON 02 MAY 2005
L12 16406 S L2
L13 0 S L12 NOT L2

FILE 'MEDLINE' ENTERED AT 16:22:54 ON 02 MAY 2005
L14 20445 S L2
L15 0 S L14 NOT L2
L16 17 S L14 AND DIFFERENT(W) CONCENTRATION?

=> s l12 and different(w)concentration?
48736 ARRAY?
12430 MICROARRAY?
797644 DNA
67604 OLIGONUCLEOTIDE?
1068128 DIFFERENT
1037046 CONCENTRATION?
9945 DIFFERENT(W) CONCENTRATION?
L17 17 L12 AND DIFFERENT(W) CONCENTRATION?

=> s l17 not l16
L18 0 L17 NOT L16